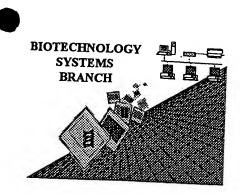
Wessendorf .

NOV 0 5 1939

RAW SEQUENCE LISTING ERROR REPORT



The Biotechnology Systems Branch of the Scientific and Technical Information Center (STIC) detected errors when processing the following CRF diskette:

Application Serial Number: 09/0/3,077

Art Unit / Team No.: 16/8

Date Processed by STIC: 8/25/9

THE ATTACHED PRINTOUT EXPLAINS THE ERRORS DETECTED.

PLEASE BE SURE TO FORWARD THIS INFORMATION TO THE APPLICANTS BY EITHER:

- 1) INCLUDING A COPY OF THIS PRINTOUT IN YOUR NEXT COMMUNICATION TO THE APPLICANTS ALONG WITH A NOTICE TO COMPLY or,
- 2) CALLING APPLICANTS AND FAXING THEM A COPY OF THE PRINTOUT WITH A NOTICE TO COMPLY

THIS WILL INSURE THAT THE NEXT SUBMISSION RECEIVED FROM THEM WILL BE ERROR FREE.

IF YOU HAVE ANY FURTHER QUESTIONS, PLEASE CALL:

MARK SPENCER 703-308-4212

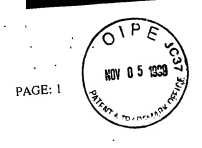


Raw Sequence Listing Error Summary

ERROR DETECTED SUGGESTED CORRECTION

SERIAL NUMBER: 09/0/3,077

AI	TN: NEW RULES CASES	S: PLEASE DISREGARD ENGLISH "ALPHA" HEADERS, WHICH WERE INSERTED BY PTO SOFTWARE
1_	Wrapped Nucleics	The number/text at the end of each line "wrapped" down to the next line.
		This may occur if your file was retrieved in a word processor after creating it.
		Please adjust your right margin to .3, as this will prevent "wrapping".
	•	revent "wrapping".
2	Wrapped Aminos	The amino acid number/text at the end of each line "wrapped" down to the next line.
		This may occur if your file was retrieved in a word processor after creating it.
		Please adjust your right margin to .3, as this will prevent "wrapping".
		wrapping.
. 3	Incorrect Line Length	The rules require that a line not exceed 72 characters in length. This includes spaces.
	-	This includes spaces.
4	Misaligned Amino Aci	d The numbering under each 5th amino acid is misaligned. This may be caused by the use of tabs
	Numbering	between the numbering. It is recommended to delete any tabs and use spacing between the numbers.
, }		to delete any tabs and use spacing between the numbers.
5 <u>U</u>	Non-ASCII	This file was not saved in ASCII (DOS) text, as required by the Sequence Rules.
		Please ensure your subsequent submission is pound in ACOUA
		Please ensure your subsequent submission is saved in ASCII text so that it can be processed.
6	Variable Length	Sequence(s) contain n's or Xaa's which represented more than one residue.
		As per the rules, each n or Xaa can only represent a single residue.
		Please present the maximum number of continuity and activities to the state of the
		Please present the maximum number of each residue having variable length and
		indicate in the (ix) feature section that some may be missing.
7	_ Patentln ver. 2.0 "bug"	A "hun" in Patentin version 2.0 has assessed the cooperate
	• 3	The state of the second of the second
		sequence(s) Normally, Patentin would automatically generate this section from the
		previously coded nucleic acid sequence. Please manually copy the relevant <220>-<223> section
		to the subsequent amino acid sequence.
8	_ Skipped Sequences	Sequence(s) micries (finterfiner)
	(OLD RULES)	Sequence(s) missing. If intentional, please use the following format for each skipped sequence: (2) INFORMATION FOR SEQ ID NO:X:
		(1) SEQUENCE CHARACTERISTICS (2)
		(i) SEQUENCE CHARACTERISTICS:(Do not insert any headings under "SEQUENCE CHARACTERISTICS")
		(x) SEGOENCE DESCRIPTION:SEQ ID NO:X:
		This sequence is intentionally skipped
		Please also adjust the "/" All INDED OF GROUP
		Please also adjust the "(iii) NUMBER OF SEQUENCES:" response to include the skipped sequence(s).
9	Skipped Sequences	Sequence(c) missing (this etc.)
	(NEW RULES)	Sequence(s) missing. If intentional, please use the following format for each skipped sequence. <210> sequence Id number
	(= : : : : : : : : : : : : : : : : : :	sequence in number
		<400> sequence id number 000
		•
10	Use of n's or Xaa's	Hen of also and less Versitations
	(NEW RULES)	Use of n's and/or Xaa's have been detected in the Sequence Listing.
	(CAT NOLES)	Use of <220> to <223> is MANDATORY if n's or Xaa's are present.
		In <220> to <223> section, please explain location of n or Xaa, and which residue n or Xaa represents.
11	Use of <213>Organism	
	(NEW RULES)	Sequence(s) are missing this mandatory field or its response.
	(NOLLS)	
12	Use of <220>Feature	Comments
	(NEW RULES)	Sequence(s) are missing the <220>Feature and associated headings.
		Use of <220> to <223> is MANDATORY if <213>ORGANISM is "Artificial" or "Unknown"
		Please explain source of genetic material in <220> to <223> section.
		(See "Federal Register," 6/01/98, Vol. 63, No. 104, pp. 29631-32) (Sec. 1.823 of new Rules)
13	Data-it	, and the state of
	Patentin ver. 2.0 "bug"	Please do not use "Copy to Disk" function of Patentin version 2.0. This causes a corrupted
		ine, resulting in missing mandatory numeric identifiers and responses (as indicated on case of the line).
		miscos, prease use "rile manager" or any other means to copy file to floppy disk.
		AKS-Biotechnology Systems Branch- 5/15/99



-->

RAW SEQUENCE LISTING PATENT APPLICATION US/09/013,077



DATE: 08/25/1999 TIME: 09:29:57

INPUT SET: S33047.raw

This Raw Listing contains the General Information Section and those Sequences containing ERRORS.

Sel item S on Even Surnew

Does Not Comply
Corrected Diskette Needed SEQUENCE LISTING 1 General Information: 2 (1) 3 (i) APPLICANT: REID, ROBERT H. 4 SADEGH-NASSERI, SCHEHERAZADE 5 6 WOLFF, MARCIA (ii) TITLE OF INVENTION: MODEL FOR TESTING IMMUNOGENICITY OF PEPTIDES 7 NAUSS, JEFFREY L. 8 9 10 (iii) NUMBER OF SEQUENCES: 15 11 12 (A) ADDRESSEE: WERTEN BELLAMY, ESQ., OFFICE OF JUDGE 13 14 ADVOCATE GENERAL, ARMY 15 (B) STREET: 901 NORTH STUART STREET 16 (C) CITY: ARLINGTON 17 18 (D) STATE: VA 3) COUNTRY: USA
F) ZIP: 22203

(V) COMPUTER READABLE FORM:
(A) MEDIUM TYPE: Floppy disk
(B) COMPUTER: IBM PC compatible
(C) OPERATING SYSTEM: PC-DOS/MS-DOS
(C) OPERATING SYSTEM: PC-DOS/MS-DOS
(D) SOFTWARE: Patentln Release #1 S. C. Wersion #1.25
(D) SOFTWARE: Patentln Release #1 S. C. Wersion #1.25
(D) NUMBER: US 08/247.884

CURRENT APPLICATION DATA

MAY-1994

OATA:

LETTON NUMBER: US 08/247.884

DATA:

LETTON NUMBER: US 08/247.884 19 (E) COUNTRY: USA 20 (F) ZIP: 22203 21 22 25 26 27 28 29 30 31 32 (viii) ATTORNEY/AGENT INFORMATION: 33 (A) NAME: BELLAMY, WERTEN F.W. 34 (B) REGISTRATION NUMEER: 27,029 35 (C) REFERENCE/DOCKET NUMBER: 252.32966PX1 36 37 (ix) TELECOMMUNICATION INFORMATION: 38 (A) TELEPHONE: 703-696-8119 39 (B) TELEFAX: 703-696-8116 40 41

42 43



PAGE: 2



RAW SEQUENCE LISTING PATENT APPLICATION US/09/013,077

DATE: 08/25/1999 TIME: 09:29:57

```
(2) INFORMATION FOR SEQ ID NO:1:
44
45
     (i) SEQUENCE CHARACTERISTICS:
46
     (A) LENGTH: 13 amino acids
 47
 48
     (B) TYPE: amino acid
     (C) STRANDEDNESS: single
 49
     (D) TOPOLOGY: linear
50
51
     (ii) MOLECULE TYPE: peptide
52
     (iii) HYPOTHETICAL: NO
 53
 54
     (iv) ANTI-SENSE: NO
 55
 56
 57
     (v) FRAGMENT TYPE: internal
     (xi) SEQUENCE DESCRIPTION: SEQ ID NO(E)
 58
 59
 60
 61
 62
     Pro Lys Tyr Val Lys Gln Asn Thr Leu Lys Leu Ala Thr
 63
 64
                     5
 65
108
     (2) INFORMATION FOR SEQ ID NO:4:
109
     (i) SEQUENCE CHARACTERISTICS:
110
     (A) LENGTH: 13 amino acids
111
     (B) TYPE: amino acid
112
113
     (C) STRANDEDNESS: single
     (D) TOPOLOGY: linear
114
115
116
     (ii) MOLECULE TYPE: peptide
     (iii) HYPOTHETICAL: NO
117
118
119
     (iv) ANTI-SENSE: NO
120
121
     (v) FRAGMENT TYPE: internal
122
123
124
     (xi) SEQUENCE DESCRIPTION: SEQ ID NO:4:
125
     Val Gly Lye Asn Ile Thr Val Thr Ala Ser Val Asp Pro
126
127
     (2) INFORMATION FOR SEQ ID NO (1):
232
                                     ) use pumered "I
233
     (i) SEQUENCE CHARACTERISTICS:
234
235
     (A) LENGTH: 13 amino acids
     (B) TYPE: amino acid
236
     (C) STRANDEDNESS: single
237
238
     (D) TOPOLOGY: linear
239
     (ii) MOLECULE TYPE: peptide
```





(C) STRANDEDNESS: single



RAW SEQUENCE LISTING PAGE: 3 PATENT APPLICATION US/09/013,077

DATE: 08/25/1999 TIME: 09:29:58

```
(iii) HYPOTHETICAL: NO
241
     (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 10:

The Ser Tyr The Pha Sand
     (iv) ANTI-SENSE: NO
242
243
244
245
246
247
248
249
250
251
347
     (2) INFORMATION FOR SEQ ID NO:14:
348
     (i) SEQUENCE CHARACTERISTICS:
349
     (A) LENGTH 82 amino acids
350
     (B) TYPE: amino acid
351
     (C) STRANDEDNESS: single
352
     (D) TOPOLOGY: linear
353
354
     (ii) MOLECULE TYPE: peptide
     (iii) HYPOTHETICAL: NO
356
357
     (iv) ANTI-SENSE: NO
358
     (v) FRAGMENT TYPE: N-terminal
359
360
361
     (xi) SEQUENCE DESCRIPTION: SEQ ID NO:14:
362
363
     Ile Lys Glu Glu His Val Ile Ile Gln Ala Glu Phe Tyr Leu Asn Pro
364
365
366
     Asp Gln Ser Gly Glu Met Phe Asp Phe Asp Gly Asp Glu Ile Phe
20 25 25 20 30 E puntury
367
368
369
     His Val Asp Met Ala Lys Lys Glu Thr Val Trp Arg Leu Glu Glu Phe
370
             35-35 40 40
371
372
     Gly Arg Phe Ala Ser Phe Glu Ala Gln Gly Ala Leu Ala Asn Ile Ala.
373
374
375
376
     Val Asp Lys Ala Asn Leu Glu Ile Met Thr Lys Arg Ser Asn Tyr Thr
                         20 %
                                           75-- 15
377
     Pro Ile
378
     80
379
380
381
     (2) INFORMATION FOR SEQ ID NO:15:
382
383
     (i) SEQUENCE CHARACTERISTICS:
384
    (A) LENGTH: 96 amino acids
385
     (B) TYPE: amino acid
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DATE: 08/25/1999 TIME: 09:29:58

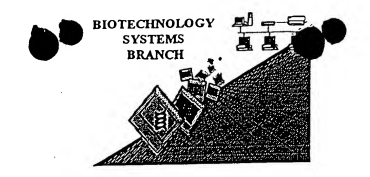
									v					IN	<i> PUT</i>	SET: S33047.raw
387	(D)	TOP	OLOGY	: 1:	inear	c										
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389	(ii)	MOI	LECUI	LE TY	PE:	pept	tide									
390	(iii	L) H	YPOTI	HETIC	CAL:	No										
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399	Glv	Asp	Thr	Ara	Pro	Ara	Phe	Leu	Trp	Gln	Leu	Lys	Phe	Glu	Cvs	His
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401	•									- •						
402	Phe	Phe	Asn	Glv	Thr	Glu	Ara	Val	Ara	Leu	Leu	Glu	Ara	Cvs	Ile	Tvr
403				20			9		25				5	30		-4-
404				20												
405	Δsn	Gln	Glu	Glu	Ser	Val	Ara	Phe	Asn	Ser	Asp	Val	Glv	Glu	Tvr	Ara
406	ASII	01	35	O_u	501		9	40		501			45		- , -	5
407			55													
408	Δla	Val	Thr	Glu	Leu	Glv	Δrσ	Pro	Asp	Δla	Glu	Tyr	Tro	Asn	Ser	Gln
409	ALU	50		014	200	017	55		шр			60				
410		70					-					• •	•			
411	T.vs	Δsn	Len	Leu	Glu	Gln	Ara	Ara	Ara	Δla	Val	Asp	Thr	Tvr	Cvs	Ara
412	65	пор	200	200	010	70	9	9	••• 9		75	F		- , -	-1-	80
413	05					, 0					. •					
414	Hie	Mot	ጥህዮ	Glv	Val	Glv	Glu	Ser	Phe	Thr	Val	Gln	Ara	Ara	Val	His
415	1113	J	ı yı	GLY	85	O _T y	OIL	DCI	1 110	90	***	04	**** 9	9	95	
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420 421																
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PAGE: 1



DATE: 08/25/1999 TIME: 09:29:59

Line	Error	Original Text
30 61 111 126 232 247 350 416	Wrong application Serial Number Wrong Sequence Number Entered (13) and Calc. Seq. Length (12) differ Wrong Amino Acid Designator Invalid Character at Sequence Number Position Wrong Sequence Number Entered (82) and Calc. Seq. Length (81) differ Wrong Amino Acid Designator	(A) APPLICATION NUMBER: Us 08/247,884 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:1: (A) LENGTH: 13 amino acids Val Gly Lye Asn Ile Thr Val Thr Ala Ser Val Asp Pro (2) INFORMATION FOR SEQ ID NO:10: (xi) SEQUENCE DESCRIPTION: SEQ ID NO:10: (A) LENGTH: 82 amino acids ??



Notice of Availability of Checker Program

Applicant Aid for Biotechnology Computer Readable Form (CRF)
Sequence Listing Submissions

The Patent and Trademark Office (PTO) has developed a computer program, called Checker, that will aid applicants in identifying and correcting errors prior to making submissions for compliance with the Requirements for Patent Applications Containing Nucleotide Sequence and/or Amino Acid Sequence Disclosures (Sequence Rules: 37CRF 1.821 through 1.825). Final rules were published in the Federal Register (55 FR18230) on May 1, 1990, and in the PTO Official Gazette (1114 Off.Gaz.PatOffice 29) on May 15, 1990.

Checker is a DOS-based software program that is intended to assist users in determining whether errors may be present in the sequence listings, and is not intended to guarantee that the submission is error-free.

The most current version of the software is available via computer downloading, details are below. Copies on diskette are also available. Updated software versions will not be automatically mailed out; any updates will be announced in the PTO Official Gazette.

The software can be accessed/requested from the following locations:

- Dial-up access through the Internet. Location is ftp://ftp.uspto.gov
 The software is in current directory: pub/checker/
 Download all the files. Cost: Free-of-charge
- 3) For diskette copies, mail to: U.S.P.T.O., OEIP, CRYSTAL PARK 3, SUITE 441 WASHINGTON DC 20231

COST FOR DISKETTE IS <u>\$ 25.00</u> METHOD OF PAYMENT:

Check payable to Commissioner of Patents and Trademarks VISA/ Mastercard/ Charge- Charges can be faxed to 703-306-2737 PTO Deposit Account

For Further Information, Contact: Arti Shah at 703-308-4212